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MOVEL METALLOPROTEASE AND DNA CODING FOR THE SAME.

② A novel metalloprotease, a DNA coding for the same, a plasmid having the sequence of the DNA, a host cell having the plasmid, and a monoclonal antibody binding specifically to the protein thereof, being useful in the medical and physiological fields for recognizing the presence of a cancer cell, diagnosing cancer malignancy, and other applications.

TECHNICAL FIELD

The present invention relates to a novel metalloproteinase useful in applications such as diagnosis of the presence of tumour cells, diagnosis of the degree of tumour malignancy, or other medical or physiological fields.

More specifically, the present invention relates to one type of metalloproteinase expressed specifically in human tumour cells and a DNA sequence encoding therefor; a plasmid having a nucleotide sequence which contains said DNA sequence; a host cell harbouring said plasmid; a method for manufacturing said protein using said host cell; a probe which hybridizes with the aforesaid DNA sequence; a method for detecting DNA or RNA containing the aforesaid sequence using said probe; and monoclonal antibodies which bind specifically to the aforesaid protein.

BACKGROUND

A group of enzymes with different substrate specificity and referred to in general as matrix metalloproteinases (hereinafter referred to as "MMPs") contributes to degradation of the extracellular matrix comprising such complex components as collagen, proteoglycan, elastin, fibronectin, and laminin.

Previously reported MMPs include interstitial collagenase (MMP-1), 72 kDa gelatinase (also known as type IV collagenase or gelatinase A; MMP-2), 92 kDa gelatinase (also known as type IV collagenase or gelatinase B; MMP-9), stromelysin-1 (MMP-3), matrilysin (MMP-7), neutrophil collagenase (MMP-8), stromelysin-2 (MMP-10) and stromelysin-3 (MMP-11).

These MMPs are a family of enzymes whose primary structure has been reported previously. With the exception of MMP-7, the primary structure among the family of reported MMPs comprises essentially an N-terminal propeptide domain, a Zn⁺ binding catalytic domain and a C-terminal hemopexin-like domain. In MMP-7 there is no hemopexin-like domain. MMP-2 and MMP-9 contain an additional gelatin-binding domain. In addition, a proline-rich domain highly homologous to a type V collagen $\alpha 2$ chain is inserted in MMP-9 between the Zn⁺ binding catalytic domain and the C-terminal hemopexin-like domain.

In highly metastatic turnour cells, there are reports of conspicuous expression of type IV collagenase (MMP-2, MMP-9) which mainly degrade type IV collagen (Cancer Res., 46:1-7, 1986; Biochem. Biophys. Res. Commun., 154:832-838, 1988; Cancer, 71:1368-1383, 1993). Likewise, it has been reported MMP-3 act as an activator of proMMP-9 (J. Biol. Chem., 267:3581-3584, 1992).

The degree of matrix metalloproteinase expression serves as an index to diagnosing the degree of cancer malignancy.

DISCLOSURE OF THE INVENTION

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The present inventors discovered a novel matrix metalloproteinase (hereinafter deferred to as "MT-MMP") and performed a structural analysis thereof.

As described hereafter, the present invention offers a novel metalloproteinase protein, DNA having a nucleotide sequence which encodes said protein, a plasmid having said DNA nucleotide sequence, a host cell harbouring said plasmid and monoclonal antibodies which specifically recognize the aforesaid metalloproteinase protein.

- 1. A protein having the amino acid sequence shown in Sequence Sheet sequence number 1.
- 2. A DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2 which encodes a protein having the amino acid sequence shown in Sequence Sheet sequence number 1.
- 3. A plasmid containing a DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2, and expressing the protein shown in Sequence Sheet sequence number 1.
- 4. A host cell harbouring a plasmid containing a DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2, and expressing the protein shown in Sequence Sheet sequence number 1.
- 5. Monoclonal antibodies which peculiarly recognize a protein having the amino acid sequence shown in Sequence Sheet sequence number 1.

The present invention is described in detail hereafter.

Using highly conserved sequences (Sequence Sheet sequence numbers 3 and 4) selected from amino acid sequences of the known matrix metalloproteinase (MMP) family, the present inventors designed and synthesized an oligonucleotide primer having the sequences denoted by Sequence Sheet sequence numbers 5 and 6. A PCR was carried out using said oligonucleotide primer and a human placental cDNA library, the PCR products obtained were sequenced, and a 390 bp DNA fragment having a sequence non-

homologous to known MMP was obtained. Using this 390 bp DNA fragment as a probe, the human placenta cDNA library was screened, and a cDNA in the positive phage clone obtained was sequenced. The nucleotide sequence is that denoted by Sequence Sheet sequence number 2. A sequence identical to the nucleotide sequence in Sequence Sheet sequence number 2 did not exist in the Genbank/EMBL DNA database, and DNA having this nucleotide sequence was ascertained to be completely novel.

The nucleotide sequence of the aforesaid cloned cDNA in Sequence Sheet sequence number 2 had a 3' non-coding sequence and open reading frame that potentially encode 582 amino acid. An initiation codon was located at nucleotide number 112, and a stop codon was present at nucleotide number 1858. It was determined that this open reading frame encoded the 582 amino acid sequence in Sequence Sheet sequence number 1, that a deduced signal sequence continued after the initiation codon, and that a hydrophobic domain (Sequence Sheet sequence number 7) specific to a membrane-binding protein of 20 or more linked hydrophobic amino acids was present from C-terminal amino acid number 533 to 562.

When homology between the amino acid sequence of MT-MMP and that of the known MMP family was analyzed, MT-MMP had high homology to the known MMP family, as shown in Figure 2. The sequences best conserved in MT-MMP were active site sequences, as well as sequences proximal to processing site between precursor and mature substance conserved in the MMP family. The fact that MT-MMP has the structural characteristics of a membrane-binding protein, and the presence in MT-MMP of a sequence of linked hydrophobic amino acids (shown in Sequence Sheet sequence number 7) not found in the rest of the MMP family, strongly suggested that MT-MMP, unlike other MMP family, is a membrane-binding MMP.

When MT-MMP expression in various human tissues was studied by Northern Blot analysis with various tissue-derived Poly(A)RNA, high expression was seen in the placenta, lung and kidney (see Figure 3). Likewise, results from Northern Blot analysis performed with RNA extracted from normal and tumour areas of human lung squamous cell carcinoma showed that MT-MMP is expressed peculiarly at tumour sites (see Figure 4).

Finally, immunoprecipitation and immunostain experiments using anti-MT-MMP monoclonal antibodies showed that the MT-MMP pertaining to the present invention is expressed on a cell membrane without secretion of a gene product, and MMP-2 activation induced by the expression of MT-MMP was observed in the cells transfected with MT-MMP gene (Nature, 370:61-65, 1994).

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Due to the achievements of the above-discussed research by the present inventors, the present invention offers a novel matrix metalloproteinase protein having the amino acid sequence in Sequence Sheet sequence number 1.

In addition, the present invention offers DNA having the nucleotide sequence in Sequence Sheet sequence number 2, which encodes a protein having the amino acid sequence in Sequence Sheet sequence number 1; a plasmid containing and capable of expressing said DNA; and a host cell harbouring said plasmid. All host cells used in general recombinant DNA technology can be used as the aforementioned host cell, including prokaryotes such as E. coli and Baci lus subtilus; eukaryotes such as yeast, COS cells, CHO cells and 3T3 cells; and insect cells such as Sf21. Expression vectors corresponding to used host cells can be used as the aforementioned plasmid.

Furthermore, the present invention offers mRNA transcribed from DNA having the nucleotide sequence in Sequence Sheet sequence number 2.

The present invention also offers a probe which hybridizes with the aforementioned DNA or RNA and specifically detects said DNA or RNA, and said probe may be one having any part of the nucleotide sequence in Sequence Sheet sequence number 2, provided said probe is labeled by a generally used radioactive isotope or enzyme or the like, hybridizes specifically with said DNA or RNA in general blotting analysis and *in situ* hybridization, and accomplishes detection.

Furthermore, the present invention offers monoclonal antibodies which bind peculiarly with the MT-MMP pertaining to the present invention.

The monoclonal antibodies pertaining to the present invention can be prepared by a well-known method such as the method of Milstein et al. (Nature, 256:495-497, 1975) using human MT-MMP as an antigen. In this method, the antigen may be native human MT-MMP, recombinant human MT-MMP, or a synthetic peptide having a partial amino acid sequence of either.

By means of the present invention, DNA having a nucleotide sequence which encodes a protein with the amino acid sequence of the novel MT-MMP pertaining to the present invention can be cloned, and such DNA and a protein encoded by such DNA can be prepared by a genetic engineering technique. Through the use of a cDNA clone of such a novel MT-MMP, techniques generally used in genetic engineering can be used to clone the aforementioned nucleotide sequence into another vector or host. Based on the aforementioned cDNA nucleotide sequence, DNA appropriately suited to a probe may be designed and prepared. In addition, based on the nucleotide sequence of the MT-MMP pertaining to the present invention,

techniques generally used in genetic engineering can be used to prepare a corresponding protein wherein appropriate mutation have been introduced into the MT-MMP amino acid sequence by substitution, deletion, insertion, displacement or addition of one or more amino acids. All such aforementioned derivatives may also be included in the present invention, provided that common metalloproteinase characteristics are conserved; namely, sequences proximal to processing site between precursor and mature substance, active site sequences and domain structure, and provided that the MT-MMP characteristic of a hydrophobic domain of linked hydrophobic amino acids present near the C terminus is conserved.

Use of the above-discussed various implementations of the present invention offers various technical means applicable to applications pertaining to diagnostic agents or diagnostic methods used for diagnosis of the presence of tumour cells or for diagnosis of the degree of tumour malignancy, as well as applications in other medical or physiological fields.

The present invention is described in detail hereafter by means of Working Examples, but the present invention is not limited by these Working Examples.

WORKING EXAMPLES

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Working Example 1 Isolation of novel metalloproteinase (MT-MMP) cDNA

(a) Construction of cDNA Library

Total RNA was extracted from human placenta tissue by a guanidine-cesium chloride method (Biochemistry, 18:5294-5299, 1979) and poly(A)⁺RNA was purified using an oligo(dT)-cellulose column. Using a purified poly(A)⁺RNA as a template and an oligo(dT) primer, cDNA was synthesized according to the Gubler-Hoffman method (Gene, 25:263-269, 1983). The ends of the cDNA were converted to blunt end with T₄ DNA polymerase, and EcoR I sites present in the cDNA were methylated by EcoR I methylase. Using T₄ DNA ligase, an EcoR I linker [d(pG-G-A-A-T-T-C-C)] and the cDNA were ligated, and cDNA possessing EcoR I sites at both ends was generated by EcoR I digestion. Using T₄ DNA ligase, this cDNA was cloned into EcoR I site of λgt11. *In vitro* packaging of this cDNA was carried out, for example, using an *in vitro* packaging kit (Amersham), and a cDNA library was thus constructed. A commercial cDNA library such as a human placenta cDNA library (Clontech) can be used as a cDNA library.

(b) Preparation of synthetic oligonucleotide primer

The sequences denoted by Sequence Sheet sequence numbers 3 (P-1) and 4 (P-2) were selected from among amino acid sequences of the known MMP family as highly conserved amino acid sequences in the MMP family, and oligodeoxynucleotide primers corresponding respectively to oligopeptide P-1 and oligopeptide P-2 were designed. Specifically, when amino acids coded by two or more codens were present in an oligopeptide, the sequences were designed as a mixture as shown in Sequence Sheet sequence numbers 5 (primer 1) and 6 (primer 2). Primer 1 and primer 2 were synthesized by a β -cyanoethyl phosphoamidite method using a DNA synthesizer (Applied Biosystems Model 392). Using a NICK column (Pharmacia) equilibrated with 10mM sodium phosphate buffer, pH 6.8 the obtained primer 1 and primer 2 were purified.

(c) Gene amplification by PCR

Using a human placenta-derived cDNA as a template and primers 1 and 2 noted in the above section (b), a PCR (PCR Technology, Stockton Press, pp. 63-67, 1989) was run.

As a result, a 390 bp PCR product was yielded. The obtained PCR product was cloned in an appropriate plasmid, e.g., pUC 119 or pBluescript, and the nucleotide sequence of the PCR product was determined using a fluorescence DNA sequencer (Applied Biosystems, Model 373A) and a Taq dye-primer cycle sequencing kit (Applied Biosystems). Among various PCR products whose nucleotide sequences were determined, PCR product A having no homology to nucleotide sequences of previously reported MMPs was obtained. PCR product A was used as a probe for screening the human placenta cDNA library noted in the foregoing section (a). ³²P labeling of the probe was generated using a random primed DNA labeling kit (Boehringer Mannhaim).

(d) Screening of novel MMP gene from cDNA library and DNA sequencing.

Host E. coli Y1090 was transfected with the human placenta cDNA library constructed in the λgt11 cited in the foregoing section (a) and plaques were formed. Specifically, Y1090 was cultured overnight in an L broth containing 0.02% maltose, and bacteria were harvested and suspended in 10mM MgSO₄. This cell suspension and a phage solution were mixed, incubated at 37°C for 15 minutes, and then the phages were adsorbed onto the host bacteria. Soft agar was added thereto, and the material was spread on an L plate (the above-noted operation is hereinafter termed "plating"). The plate was incubated overnight at 42 °C and a plaque was formed, after which a nylon filter (e.g., Hibond-N, Amersham) or a nitrocellulose filter (e.g., HATF, Millipore) was placed onto the plate and left in place for approximately 30 seconds. The filter was gently peeled and immersed in an alkaline denaturant (0.1M NaOH, 1.5M NaCl) for 30 seconds, then immersed in a neutralizing solution (0.5M Tris-HCl buffer, pH 8 containing 1.5M NaCl) for 5 minutes. The filter was then washed with 2x SSPE (0.36M NaCl, 20mM NaH2PO4, 2mM EDTA) and dried. The foregoing plaque-to-filter transfer was repeated, and at least two filters were prepared. However, plate contact time for the second and subsequent filters was extended to approximately 2 minutes. Filters were baked 2 hours at 80 °C and DNA was thus fixed. The two filters, at a minimum, prepared from one plate were respectively washed 1 hour at 42 °C in a wash solution (50mM Tris-HCl buffer, pH 8.0 containing 1M NaCl, 1mM EDTA and 0.1% SDS), placed in a hybridization bag, and prehybridization was carried out by 6 to 8 hours immersion at 42°C in a prehybridization solution [50% formamide, 5x Denhardt's solution (0.2% bovine serum albumin, 0.2% polyvinylpyrolidone), 5x SSPE, 0.1% SDS, 100µg/ml heat-denatured salmon sperm DNA]. Next, the ³²P-labeled probe noted in section (c), heat-denatured for 5 minutes at 100 °C, was added to the prehybridization solution, and hybridization was carried out overnight at 42 °C. After hybridization was complete, the filters were washed at room temperature with an excess of 2x SSC solution containing 0.1% SDS. Next, the filters were placed for 15 minutes at 68 °C in 1x SSC solution containing 0.1% SDS. The filters were then dried, layered with X-ray film (Kodak XR), and 1 week autoradiography was then carried out at -70 °C. The X-ray films were developed, replica filters in duplicate produced from one plate were piled up each other, and signals that appeared precisely same place on duplicate filters were marked. Plaques corresponding to marked signals were suspended in SM solution (50mM Tris-HCl buffer, pH 7.5 containing 0.1M NaCl and 10mM MgSO₄). These phage suspensions were appropriately diluted and plating was performed, screening similar to that noted above was carried out, and recombinant phages were obtained.

(e) Preparation of recombinant λgt11 DNA

Each cloned phages was plated, incubated for 3 hours at 42 °C, and incubated overnight at 37 °C. Several drops of chloroform was then added to the SM solution and the material was left at room temperature for 30 minutes. The SM solution together with the upper layer of soft agar was then scraped off, and centrifuged. Polyethylene glycol was added to a 10% final concentration in the supernatant, and the material was mixed and left at 4 °C for 1 hour. The material was then centrifuged, the supernatant was discarded, and phage particles were collected. The phage particles were suspended in SM solution and purified by a glycerol gradient ultracentrifugation method (see "Molecular Cloning, a Laboratory Manual", T. Maniastis et al., Cold Spring Harbor: Laboratory Press pp. 2.78, 1989). The phages obtained were suspended in SM solution and treated with DNase I and RNase A. A mixture of 20mM EDTA, 50µg/ml proteinase K, and 0.5% SDS was then added, and the material was incubated at 65 °C for 1 hour. The material was then subjected to phenol extraction and diethylether extraction, and DNA was precipitated by ethanol precipitation. The DNA obtained was washed with 70% ethanol, dried, and dissolved in TE solution (10mM Tris-HCl buffer, pH 8 containing 10mM EDTA).

(f) Sequencing of the insertion fragment

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The \(\lambda\)gt11 DNA prepared in the above section (e) was digested with EcoR I, an insertion fragment was excised and purified, and cloned into the EcoR I site of a pBluescript (Stratagene) vector. E. coli NM522 XLI-Blue was transformed with this recombinant pBluescript. The F' transformed cells were selected, infected with helper phage VCSM13 (Stratagene), and cultured overnight. The culture was centrifuged and the bacteria were removed, and PEG/NaCI was added to precipitate the phag s. The precipitate was suspended in TE solution, and single-stranded DNA was extracted with phenol and recovered by ethanol precipitation. The single-stranded DNA was sequenced using a fluorescence DNA sequencer (Applied Biosystems, Model 373A) and a Taq dye-primer cycle sequencing kit (Applied Biosystems). The total length

of the sequence determined was 3403 base pairs, and the sequence thereof is denoted by Sequence Sheet sequence number 2. The nucleotide sequence in Sequence Sheet sequence number 2 was searched using the Genbank/EMBL DNA database, but an identical sequence did not exist.

(g) Analysis of Gene Product

Hydrophilic and hydrophobic values of the amino acid sequence denoted by Sequence Sheet sequence number 1, as predicted from the nucleotide sequence denoted by Sequence Sheet sequence number 2, were calculated by the Kyte-Doolittle method (J. Mol. Biol., 157:105-132, 1982), and the hydrophilic and hydrophobic distribution plot shown in Figure 1 was determined. A hydrophobic domain comprising a sequence of 20 or more linked hydrophobic amino acids peculiar to a membrane binding protein was present from position 533 to position 562 of the C-terminal region of Sequence Sheet sequence number 1, and the sequence thereof is shown in Sequence Sheet sequence number 7. Such a sequence of linked hydrophobic amino acids does not exist in previously known MMPs.

When the homology of the amino acid sequence in Sequence Sheet sequence number 1 was compared to reported MMPs amino acid sequences, the amino acid sequence in Sequence Sheet sequence number 1 showed homology with the MMP family. Specifically, processing site between precursor and active enzyme and active site conserved to an extremely high degree among MMP family were each highly conserved in MT-MMP as well (Sequence Sheet sequence number 1, amino acids numbers 88-97 and 112-222).

Working Example 2 Gene Expression

(a) Expression in Tissues

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Using ³²P-labeled PCR product A noted in Working Example 1, section (c) as a probe, hybridization was performed with poly(A)⁺ RNA blotted membrane, human multiple tissue Northern Blots (Clontech), which contains poly(A)⁺ RNA from human heart, brain, placenta, lung, liver, skeletal muscle, kidney and pancreas. Human multiple tissue Northern Blot filters wetted with 3x SSC (0.45M NaCl, 0.045M trisodium citrate•2H₂0, pH 7.0) were prehybridized for 2 to 3 hours in a prehybridization solution (0.75M NaCl, 2.5mM EDTA, 0.5x Denhardt's solution, 50% formamide and 20mM Tris-HCl buffer, pH 7.5 containing 1% SDS) with gentle agitation. Next, a heat-denatured probe was added to the hybridization solution (10% sodium dextran and 50µg/ml denatured salmon sperm DNA-containing prehybridization solution), the prehybridization solution was replaced, and hybridization was performed overnight at 43°C. After hybridization was complete, the filters were washed with 2x SSC containing 0.1% SDS. Next, the filters were placed for 15 minutes at 68°C in 1x SSC containing 0.1% SDS. The filters were then dried, layered with X-ray film (Kodak XR), and 1 week autoradiography was then carried out at -70°C. The size of the MT-MMP gene transcripts was 4.8 kb in each tissue. When the developed X-ray films were traced by a densitometer and signal intensity was measured, among the investigated tissues, MT-MMP genes were found to be highly expressed in the lung, placenta and kidney.

(b) Expression in Tumour Tissues

Normal and tumour tissues were taken from samples of two squamous cell carcinomas human lung, respectively, and total RNA was extracted by a guanidine-cesium chloride method. 10µg of each said RNA was applied to 1% agarose electrophoresis and then transferred onto a nylon membrane. Hybridization was then carried out with the ³²P-labeled probe noted in Working Example 1, section (c). Hybridization and autoradiography tracing were performed as described in the foregoing section (a). In each human lung squamous cell carcinoma, significantly higher expression were seen in tumour tissue (see Figure 4 T) than in normal tissue (see Figure 4 N).

Working Example 3 Preparation of Monoclonal Antibodies

(a) Preparation of Polypeptides as Antigen

From the MT-MMP amino acid sequence denoted by Sequence Sheet sequence number 1, sequences denoted by Sequence Sheet sequence numbers 8, 9 and 10 (sequence of Sequence Sheet sequence number 1 amino acid numbers 160-173, 320-333, and 498-512, respectively; hereinafter termed polypeptide

A, polypeptide B and polypeptide C, respectively) were selected as specific sequences having low homology to other members of MMP family. These polypeptides were synthesized by Fmoc-BOP method using a peptide synthesizer (MilliGen/Biosearch, Peptide Synthesizer 9600), and cysteine was introduced at the N-terminus. Each synthesized peptide was purified by high speed liquid chromatography.

(b) Preparation of Each Polypeptides and Keyhole Limpet Hemocyanin Complexes

2 mg of keyhole limpet hemocyanin (KLH) dissolved in 1 ml of 0.1M phosphate buffer, pH 7.5 and 1.85 mg N-(ε-maleimidocaproyloxy)succinimide dissolved in 200 μl dimethylformamide were mixed and incubated at 30 °C for 30 minutes. Next, the above-noted mixture was applied to gel filtration by PD-10 (Pharmacia) equilibrated with 0.1M phosphate buffer, pH 7.0. KLH-bound maleimide was collected and concentrated to less than 1.5 ml. Each polypeptide synthesized in the foregoing section (a) was respectively dissolved in 1 ml of 0.1M phosphate buffer, pH 7.0 and mixed with KLH-bound maleimide at a molar ratio representing a factor of 50. This material was then incubated 20 hours at 4 °C, and KLH-polypeptide complexes were thus prepared.

(c) Preparation of Antibody-producing Cells

As an initial immunization, eight-week-old Balb/c female mice were given an intraperitoneal administration of 250 µg of a complex of KLH and, respectively, polypeptide A, polypeptide B or polypeptide C prepared in the above section (b), in Freund's complete adjuvant. After 18 days, the respectively immunized mice were boosted intraperitoneally with 200 µg of the respective complexes dissolved in 0.1M phosphate buffer, pH 7.5. After 32 days, a final immunization of 100 µg of each complex was administered intravenously as the booster immunization. Three days thereafter, splencytes were extirpated and splencyte suspensions were prepared.

(d) Cell Fusion

Fusion with 8-azaguanine-resistant myeloma cell SP2 (SP2/O-Ag14) was performed according to a modifying method of Oi et al (see Selected Methods in Cellular Immunology, Mishell, B.B. and Shiigi, S. M., ed., W.H. Freeman and Company pp. 351-372, 1980). Fusion of myeloma cell SP2 with karyo-splencytes from mice immunized with the polypeptide A-KLH complex is discussed in details, hereafter.

Through the following procedures, karyo-splencytes prepared in the foregoing section (c) (cell viability 100%) were fused in a 5:1 ratio with myeloma cells (cell viability 100%). A polypeptide A-immunized splencyte suspension and myeloma cells were separately washed in RPMI 1640 medium. The material was then suspended in the same medium, and 3x108 cells of karyo-splencytes and 6x107 cells of myeloma cells were mixed for fusion. The cells were then precipitated by centrifugation, and all the supernatant was completely discarded by suction. 2.0 ml of PEG 4000 solution (RPMI 1640 medium containing 50% [w/v] polyethylene glycol 4000) prewarmed at 37 °C was added dropwise to the precipitated cells over 1 minute, 1 minute stirring was performed, and the cells were resuspended and dispersed. Next, 2.0 ml of RPMI 1640 medium prewarmed at 37°C was added in a dropwise fashion over 1 minute. After repeating the same operation once more, 14 ml of RPMI 1640 medium was added dropwise over 2 to 3 minutes under constant stirring, and the cells were dispersed. The dispersion was centrifuged and the supernatant was completely discarded by suction. Next, 30 ml of NS-1 medium (RPMI 1640 medium containing filter-sterilized 15% [w/v] fetal calf serum [JRH Biosciences]) prewarmed at 37°C was rapidly added to the precipitated cells, and the large cell clumps were carefully dispersed by pipetting. The dispersion was then diluted by adding 30 ml of NS-1 medium, and 6.0x105 cells/0.1 ml/well was added to a polystyrene 96-microwell plate. The above-noted cell-filled microwells were cultured in 7% carbonic acid gas/93% atmospheric air at 37 °C and 100% humidity.

In the case of splencytes derived from mice immunized with the polypeptide B-KLH complex, 6.4x10⁸ cells of splencytes and 1.28x10⁸ cells of myeloma cells were mixed, and respectively, 4.3 ml, 38.7 ml and 129 ml of the above-used PEG 4000 solution, RPMI 1640 medium and NS-1 medium were used. In the case of splencytes derived from mice immunized with the polypeptide C-KLH complex, 6.8x10⁸ cells of splencytes and 1.36x10⁸ cells of myeloma cells were mixed, and 4.5 ml, 40.5 ml and 135 ml of respectively PEG 4000 solution, RPMI 1640 medium and NS-1 medium were used.

(e) Selective Amplification of Hybridomas by Selective Culture Medium

On the day following the start of culturing mentioned in the above section (d) (Day 1), 2 drops (approx. 0.1 ml) HAT culture medium (100 µM hypoxanthine, 0.4 µM aminopterin and 16 µM thymidine added to NS-1 culture medium) were added to the cells with a Pasteur pipette. On Days 2, 3, 5 and 8, half of each culture medium (approx. 0.1 ml) was replaced with fresh HAT medium, and on Day 11, half of each culture medium was replaced with fresh HT culture medium (HAT culture medium not containing aminopterin). On Day 14, for all the wells in which hybridoma growth was observed to the naked eye, positive wells were investigated by enzyme-linked immunoadsorbent assay (ELISA). Specifically, the polystyrene 96-well plate was respectively coated with polypeptides A, B and C serving as antigens, washed using PBS for washing (containing 0.05% Tween 20), and unadsorbed peptides were thus removed. In addition, the uncoated portion of each well was blocked with 1% BSA. 0.1 ml of supernatant from wells in which hybridoma growth was confirmed was added to each polypeptide-coated well, and the plate was stood at room temperature for approximately 1 hour.

Horseradish peroxidase-labeled goat anti-mouse immunoglobulin was added as a secondary antibody, and the plate was again stood at room temperature for approximately another 1 hour. A substrate of hydrogen peroxide and o-phenylenediamine was added, and the degree of color development was measured as absorbance at 492 nm using a microplate light absorbency measuring device (MRP-A4, Tosoh).

(f) Hybridoma Cloning

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Hybridomas in wells positive with respect to individual antigen peptides, as obtained in the foregoing section (e), were monocloned according to the limiting dilution method. Specifically, hybridomas were diluted to 5, 1 and 0.5 per well and were respectively added to 36, 36 and 24 wells of a 96 microwells. On Day 5 and Day 12, approximately 0.1 ml NS-1 medium was added to each well. Approximately 2 weeks after cloning began, the ELISA noted in section (e) was performed for groups in which sufficient hybridoma growth was visually confirmed and 50% or more wells were negative for colony formation. If all tested wells were not positive, 4 to 6 antibody-positive wells in which the number of colonies was 1 were selected, and recloning was performed. Finally, as shown in Table 1 and Table 2, 12, 20 and 9 hybridomas were obtained which produced monoclonal antibodies against polypeptide A, polypeptide B or polypeptide C, respectively.

(g) Hybridoma Culturing and Monoclonal Antibody Purification

Each obtained hybridoma was cultured in NS-1 medium and a 10 to 100 μg/ml concentration of monoclonal antibody was successfully obtained from the supernatant thereof. In addition, BALB/c mice given an one week prior intraperitoneal administration of pristane were given a similar intraperitoneal administration of 1x10⁷ cells of obtained hybridomas, and after 1 to 2 weeks, abdominal fluid containing 4 to 7 mg/ml of monoclonal antibody was successfully obtained. The abdominal fluid obtained was salted out by 40% saturated ammonium sulfate, and IgG class antibodies were adsorbed to Protein A Affigel (Bio-Rad) and purified by elution with a 0.1M citric acid buffer, pH 5.

(h) Determination of Monoclonal Antibody Class and Subclass

In accordance with the above-discussed ELISA, the supernatant of monoclones obtained in section (f) were added to microtitration plates respectively coated with polypeptide A, polypeptide B or polypeptide C. After washing with PBS, isotype-specific rabbit anti-mouse IgG antibodies (Zymed Lab.) were added. After washing with PBS, horseradish peroxidase-labeled goat anti-rabbit IgG (H+L) was added, and class and subclass were determined using hydrogen peroxide and 2.2'-azino-di(3-ethylbenzthiazolinic acid) as a substrate.

(i) Specificity of Anti-MT-MMP Monoclonal Antibodies

The cross-reactivity of five varieties of anti-MT-MMP monoclonal antibodies (monoclone numbers 113-5B7, 113-15E7, 114-1F1, 114-2F2 and 118-3B1) exhibiting a positive reaction against a human MT-MMP peptide was determined by the ELISA noted in the foregoing section (e), using as respective antigens: proMMP-1 (Clin. Chim. Acta, 219:1-14, 1993), proMMP-2 (Clin. Chim. Acta, 221:91-103, 1993) and proMMP-3 (Clin. Chim. Acta, 211:59-72, 1992) respectively purified from the supernatant of nomal human skin

fibroblast (NB1RGB) culture; proMMP-7 purified from the supernatant of human rectal carcinoma cell (CaR-1) culture (Cancer Res., 50:7758-7764, 1990), proMMP-8 purified from human neutrophils (Biol. Chem. Hoppe-Seyler, 371 supp:295-304, 1990) and proMMP-9 purified from the supernatant of human fibrosarcoma cells (HT1080) culture (J. Biol. Chem., 267: 21712-21719, 1992).

Specifically, using a polystyrene 96-well plate, each well was coated by adding 50 ng/well of purified MMP-1, MMP-2, MMP-3, MMP-7, MMP-8 and MMP-9, respectively. Washing was performed with PBS for washing and non-adsorbed antigen was removed, and the uncoated portion of each well was blocked with PBS containing 3% skim milk. 1 µg/well of each MT-MMP monoclonal antibody was respectively added to each well and stood at room temperature for approximately 1 hour. After washing plate, peroxidase-labeled goat anti-mouse immunoglobulin was added as a secondary antibody, and the plate was again stood at room temperature for approximately 1 hour. A substrate of hydrogen peroxide and o-phenylene diamine was added, and the degree of color development was measured absorbance at 492 nm using a microplate light absorbency measuring device (MRP-A4, Tosoh).

In results, as shown in Table 3, each anti-MT-MMP monoclonal antibody showed no reactivity against purified MMPs other than the MT-MMP supplied for testing.

TABLE 1

Polypeptide	Monoclone No.	Subclass/Chain
Α	114-1F2	γ1/x
	114-2F2	γ1/x
	114-3H7	· γ1/x
	114-5E4	γ1/x
	114-6G6	γ1/x
	114-8D10	γ1/x
	114-9H3	μ/x
,	114-15E8	γ1/x
	114-16C11	γ1/x
	114-18E4	γ1/x
	114-19F11	γ1/x
	114-20H5	μ/x
В	113-1E3	γ3/x
	113-2E9	γ3/x
	113-3F6	γ2b/x
	113-4H7	γ3/x
·	113-5B7	γ3/x
	113-7C6	γ2b/x
	113-9G9	γ3/x
•	113-10F2	γ3/x
	113-13G11	γ3/x
	113-15E7	γ3/x
	113-16H8	γ3/x
	113-17G12	μ/x
	113-19A10	μ/x ·
	113-20G11	γ3/x
	113-21H3	γ1/x
	113-26D3	μ/x
	113-44C1	γ1/x
	113-46B7	γ1/x
	113-53G5	μ/x
	113-63E8	γ1/x

TABLE 2

Polypeptide	Monoclone No.	Subclass/Chain		
С	118-3B1	√2b/x		
Ū	118-6F3	γ2b/x		
	118-8D11	γ1/x		
	118-9B11	γ1/x		
	118-13D11	α/x		
	118-18C12	γ1/x		
	118-20A3	γ2b/x		
	118-25C3	γ1/x		
	118-26F5	· γ3/x		

TABLE 3

Monoclone No.	Cross reactivity										
	MMP-1	MMP-2	ммр-3	MMP-7	MMP-8	ммр-9					
113-5B7	-	•	-	-	-	-					
113-15E7	-		-	-	-	-					
114-1F2	-	-		-	•	•					
114-2F2	-		-	-	-	-					
118-3B1	-	-	-	-	-	-					

Working Example 4 Expression and Identification of Gene Product

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By means of EcoR I cleavage, an insertion fragment was excised from the recombinant pBluescript containing a cloned MT-MMP gene, constructed in section (f) of Working Example 1. Cloning was then carried out at an EcoR I site of the eukaryotic expression vector pSG5 (Stratagene). Then, human fibrosarcoma cells (HT1080) were transfected with said recombinant pSG5 by a calcium phosphate method. Specifically, 20 μg of recombinant pSG5 and 62 μI of 2M CaCl₂ was added to distilled water, and 2x HBSP solution (50mM HEPES buffer, pH 7.1 containing 1.5mM Na₂HPO₄, 10mM KCl, 280mM NaCl and 12mM glucose) was added to the bottom of the tube to form a total volume of 1 ml. This material was mixed, stood at room temperature for approximately 30 minutes, and thorough precipitate formation was carried out. The precipitate was dispersed by pipetting, added dropwise to HT1080 cells and incubated for approximately 4 hours in a CO₂ incubator. Next, the culture medium was removed, a 15% glycerol solution was added and treated for 1 to 3 hours, the glycerol was discarded by suction, washed with PBS and fresh culture medium containing ³⁵S-methionine was added. Culturing was continued, and cellular proteins were labeled by ³⁵S. Incidentally, expression of MT-MMP genes in HT1080 cells cannot be detected by Northern Blot analysis.

The cells were incubated for 1 hour at 4°C in a lysing buffer solution (0.01M Tris-HCl buffer, pH 8 containing 1% Triton X-100, 1% bovine hemoglobin, 1mM iodoacetamide, 0.2U trypsin inhibitor, 1mM PMSF and 0.14M NaCl). The cell lysate was centrifuged and the supernatant was recovered. Sepharose-4B (Pharmacia) coupled with a monoclonal antibody obtained in Working Example 3 was added to the supernatant, the material was incubated at 4°C for 2 hours with agitation, and immunoprecipitation was carried out. Monoclonal antibodies against polypeptide A used in immunoprecipitation were two of the 12 obtained in Working Example 3 which had low non-specific reactivity (monoclone numbers 114-1F2 and 114-2F2 [Assignment No. FERM BP-4743]). Next, Sepharose 4B coupled with monoclonal antibodies subjected to immunoprecipitation were precipitated by centrifugation, washed three times with a washing solution (0.01M Tris-HCl buffer, pH 8 containing 1% Triton X-100, 1% bovine hemoglobin and 0.14M NaCl), and lastly, washed with 0.05M Tris-HCl buffer, pH 6.8. A sample buffer for SDS polyacrylamide electrophoresis was added to washed Sepharose-4B coupled with a monoclonal antibody, boiled 5 minutes at

100 °C, and SDS polyacrylamide electrophoresis was carried out. The electrophoresed gel was layered with X-ray film (Kodak XR), 1 week autoradiography was then carried out at -70 °C, and the developed X-ray films were traced by a densitometer to measure signal intensity. With each of the anti-MT-MMP monoclonal antibodies used (monoclone numbers 114-1F2 and 114-2F2), the immunoprecipitate contained a 63 kDa protein. In cells transfected with a pSG5 vector alone not containing an MT-MMP gene as a control, anti-MT-MMP monoclonal antibodies (monoclone numbers 114-1F2 and 114-2F2) did not precipitate a 63 kDa protein. The 63 kDa molecular weight of the protein detected by immunoprecipitation nearly matched a molecular weight of 65.78 kDa calculated from the amino acid sequence denoted by Sequence Sheet sequence number 1. In addition, a variant MT-MMP expression plasmid was constructed in which amino acids from position 13 to position 101 were deleted from the amino acid sequence denoted by Sequence Sheet sequence number 1, HT1080 cells was transfected with said variant as stated above, and immunoprecipitation was carried out. With HT1080 cells to which the variant MT-MMP gene was introduced, a 63 kDa protein was not detected, and a 55 kDa protein was detected. This molecular weight matched a molecular weight predicted from the introduced deletion.

EXPERIMENTAL EXAMPLE

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(a) Activation of proMMP-2 by MT-MMP Expression

Recombinant pSG5 carrying a cloned MT-MMP gene, constructed in Working Example 4, and a pSG5 vector alone, serving as a control, similarly transfected into HT1080 cells by the calcium phosphate method mentioned in Working Example 4, or into mouse embryonic fibroblasts NIH3T3. However, a regular fresh culture medium was used in lieu of the fresh culture medium containing ³⁵S-methionine. Both the HT1080 cells and the NIH3T3 cells secreted proMMP-2 and proMMP-9 (corresponding respectively to the 66 kDa and 97.4 kDa bands in Figure 6), and in cells transfected with an MT-MMP gene, MT-MMP expression was confirmed by immunoprecipitation experiments (See Working Example 4).

The transfectants obtained were cultured for 24 hours in a serum free medium and the recovered culture supernatant was supplied for zymography. The culture supernatant was mixed with an SDS polyacrylamide electrophoresis buffer (non-reducing condition) and left at 4°C overnight. Electrophoresis was then performed at 4°C, with a 20 mA current, using a 10% polyacrylamide gel containing 1 mg/ml casein. After electrophoresis, the gel was washed with a gelatinase-buffer (Tris-HCl buffer, pH 7.6 containing 5mM CaCl₂ and 1 μ M ZnSO₄) containing 2.5% Triton X-100 with gentle agitation for 15 minutes, and this operation was repeated twice. Next, the gel was immersed in a gelatinase-buffer containing 1% Triton X-100 and stood at 37°C overnight. The buffer was discarded and the gel was stained for 1 hour with 0.02% Coomassie Brilliant Blue-R (dissolved in 50% methanol/10% acetic acid) and destained by immersion in a destaining solution (5% methanol, 7.5% acetic acid).

As shown in Figure 6, MT-MMP gene-transfected HT1080 cells produced new 64 kDa and 62 kDa bands, confirming proMMP-2 activation. This active-form MMP-2 exhibited the same molecular weight as an active-form MMP-2 molecule induced by treatment of cells with 100 µg/ml of concanavalin A and reacted specifically against anti-MMP-2 monoclonal antibodies. This activation was not observed in a control transfected with a vector alone. Likewise, proMMP-9 showed no change in molecular weight and no activation similar to that observed in control cells. Such activation of proMMP-2 depending on MT-MMP expression was also observed in MT-MMP gene-transfected NIH3T3 cells.

(b) Activation of ProMMP-2 by MT-MMP Expression Cell Membrane Fraction

In a manner similar to that noted in the above section (a), African green monkey kidney-derived COS-1 cells were transfected with recombinant pSG5 containing cloned MT-MMP gene, or with control pSG5 vector alone by a calcium phosphate method. A cell membrane fraction was then prepared from the obtained transfectant according to the method of Strongin et al. (J. Biol. Chem., 268:14033-14039, 1993).

The transfectant was washed with PBS, and cells were harvested by centrifugation and suspended in a 25mM Tris-HCl buffer, pH 7.4 containing 8.5% sucrose, 50mM NaCl, 10mM N-ethylmaleimide, 10 µg/ml aprotinin, 1 µg/ml pepstatin A, 1 µg/ml leupeptin and 1mM phenylmethylsulfonyl fluoride. The cell suspension was homogenized in a Dounce homogeniz r, and the homogenate was centrifuged (3000x g, 10 min., 4°C). The r sulting supernatant was ultracentrifuged (100,000x g, 2 hours) and the precipitate was suspended in a 25mM Tris-HCl buffer, pH 7.4 containing 50mM NaCl, 10mM N-ethylmaleimide, 10 µg/ml aprotinin, 1 µg/ml pepstatin A, 1 µg/ml leupeptin and 1mM phenylmethylsulfonyl fluoride. This suspension was fractionated by discontinuous sucrose density gradient centrifugation (20, 30, 50, 60% sucrose

solutions; 100,000x g; 2 hours; 4°C), and bands of cell membrane fractions appeared were recovered. These fractions were precipitated again by ultracentrifugation (100,000x g, 2 hours), suspended in 25mM HEPES/KOH buffer, pH 7.5 containing 0.1mM CaCl₂ and 0.25% Triton X-100, and adjusted to a final protein concentration of 1-2 mg/ml. This suspension was ultracentrifuged (100,000x g, 1.5 hours, 4°C) to remove insoluble residue, and the supernatant obtained was taken as a cell membrane fraction.

Cell membrane fractions (protein content 20 µg) respectively prepared from untreated COS-1 cells or from COS-1 cells transfected with pSG5 vector alone or pSG5 vector with an MT-MMP gene were incubated with HT1080 cell culture supernatant at 37 °C for 2 hours. Using these samples, the zymography noted in the above section (a) was performed.

In the results, new 64 kDa and 62 kDa bands appeared and the activation of proMMP-2 present in HT1080 cell culture supernatant was observed only when cell membrane fractions derived from MT-MMP gene-transfected COS-1 cells were used (see Figure 7), and the activation of proMMP-2 was inhibited by the addition of recombinant (r) human TIMP-2. These results exhibited the activation of proMMP-2 by MT-MMP expressed on a cell membrane.

(c) Stimulation of cellular invasion in vitro due to MT-MMP expression

Invasion of cells was assayed by modified Boyden Chamber method (Cancer Res., 47:3239-3245, 1987), and operations were carried out in accordance with the manufucture's instructions for a Biocoat Matrigel Invasion Chamber (Becton Dickinson).

In a manner similar to that noted in the foregoing section (a), HT1080 cells or NIH3T3 cells were transfected with recombinant pSG5 carrying a cloned MT-MMP gene, or a control pSG5 vector alone, by a calcium phosphate method, and each of these host cells secreted proMMP-2. The resulting transfectants were then suspended in DMEM medium containing 0.1% BSA, and 2x10⁵ cells were seeded onto an uncoated filter (pore size 8 µm) or a preswelled Matrigel Coat filter in a Biocoat Matrigel Invasion Chambers. After 24 hours incubation in a CO₂ incubator at 37 °C, the filters were fixed by 10 seconds immersion in methanol. The filters were then stained by hematoxylin for 3 minutes, washed, and stained by eosin for 10 seconds, and the number of cells invaded the bottom surface of the filters were counted under a light microscope (at a magnification of x 400).

In the MT-MMP gene-transfected HT1080 cells and NIH3T3 cells, more than twice as many invading cells were seen compared to cells transfected with the control vector alone (See Figure 8 Matrigel). Specifically, MT-MMP expression was seen to stimulate cellular invasion. Furthermore, the addition of 10 µg/ml of r-human TIMP-2 to this assay system clearly suppressed cellular invasion (see Figure 8 Matrigel+r-human TIMP-2).

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 shows hydrophilic and hydrophobic distribution diagrams for the amino acid sequence of MT-MMP, according to the Kyte-Doolittle method.

Figures 2A, 2B, 2C, 2D, 2E, 2F, 2G and 2H are figures comparing sequential homology between the amino acid sequences of MT-MMP and those of the known MMP family (MMP-1, MMP-2, MMP-3, MMP-7, MMP-8, MMP-9, MMP-10 and MMP-11). Letters in each figure indicate respective amino acids; A corresponding to Ala, C to Cys, D to Asp, E to Glu, F to Phe, G to Gly, H to His, I to Ile, K to Lys, L to Leu, M to Met, N to Asn, P to Pro, Q to Gln, R to Arg, S to Ser, T to Thr, V to Val, W to Trp and Y to Tyr. Figures 2A through 2H are an integral unit and comprise a single figure.

Figure 3 shows a relative expression of MT-MMP mRNA in various human tissues, according to Northern blot analysis.

Figure 4 shows a relative expression of MT-MMP mRNA in a normal tissue and a tumour tissue of two samples of human lung squamous cell carcinoma, according to Northern blot analysis.

Figure 5 shows results for detection, by immunoprecipitation, of MT-MMP proteins expressed in HT1080 cells transfected with MT-MMP cDNA. The figure shows a scan by a densitometer, and the darkened areas indicate the location of MT-MMP immunoprecipitated by anti-MT-MMP monoclonal anti-body.

Figure 6 shows an activation of proMMP-2 by expression of MT-MMP, according to zymography of culture supernatant from HT1080 and NIH3T3 cells transfected with MT-MMP cDNA.

Figure 7 shows an activation of proMMP-2 by a cell membrane fraction of COS-1 cells transfected with MT-MMP cDNA, according to zymography.

Figure 8 shows a stimulation of the cellular invasion by expression of MT-MMP, according to a partially modified Boyden chamber method.

5	[Sequence Sheet 1]										
	Sequence No.: 1										
	Length of sequence: 582										
10	Type of sequence: Amino acid										
	Topology: Linear										
	Class of sequence: Protein										
15	Sequence										
	Net Ser Pro Ala Pro Arg Pro Ser Arg Cys Leu Leu Leu Pro Leu										
	1 5 10 15										
20	Leu Thr Leu Gly Thr Ala Leu Ala Ser Leu Gly Ser Ala Gln Ser										
	20 25 30										
05	Ser Ser Phe Ser Pro Glu Ala Trp Leu Gln Gln Tyr Gly Tyr Leu										
25	35 40 45										
•	Pro Pro Gly Asp Leu Ars Thr His Thr Gln Ars Ser Pro Gln Ser										
30	50. 55 60										
	Leu Ser Ala Ala ile Ala Ala Met Gin Lys Phe Tyr Gly Leu Gin										
	65 70 75										
35	Val The Gly Lys Ala Asp Ala Asp The Net Lys Ala Net Arg Arg										
	80 85 90										
	Pro Arg Cys Cly Val Pro Asp Lys Phe Gly Ala Glu ile Lys Ala										
40	95 100 105										
	Asn Val Arg Arg Lys Arg Tyr Ala lie Gin Gly Leu Lys Trp Gin										
	110 115 120										
45	His Asn Glu lie Thr Phe Cys lie Gin Asn Tyr Thr Pro Lys Yal										
	125 130 135										
50	Gly Glu Tyr Ala Thr Tyr Glu Ala lle Arg Lys Ala Phe Arg Val										
50	140 145 150										
	Trp Glu Ser Ala Thr Pro Leu Arz Phe Arz Glu Val Pr Tyr Ala										
55	155 160 165										

	[Se	que	nce	She	et	2]									
	Seq	uen	ce N	10.:	1	(cor	ntin	ued) .					÷	
5	Tyr	He	Arz	Clu	Cly	His	Glu	irs	Gin	Ala	ASP	He	Hèt	He	Phé
					170					175					180
	Phe	Ala	Clu	Cly	Phe	His	Gly	ÅSP	Års	The	Ala	Phe	ASP	Cly	Glu
0					185					190			•		195
	Gly	Gly	Phe	Leu	Ada	His	λla	Tyr	Phe	Pro	Cly	Pro	Åsn	ile	Gly
					200					205		•			210
5	Gly	Asp	The	His	Phe	ÅSP	Ser	Ala	Glu	Pro	Trp	Thr	Yal	Arg	۱sn
					2.15		•			220	•		,		225
	Glu	ASP	Leu	Asn	Gly	Asin	l sp	He	Phe	Leu	Ya i	Ala	Yal	His	Glu
20					230				٠.	235					240
	Leu	Gly	His	Ala	Leu	G:1 y	Leu	Clu	His	Ser	Ser	ÅSP	Pro	Ser	ila
					245					250					255
25	lle	Hel	Ala	Pro	Phe	Tyr	Gln	Trp	Hel	ķsp.	Thr	Glu	Lys	Phe	Yal
					260			•		265					270
	Leu	Pro	His	Ţyr	ÅSP	Pro	årg	Gly	He	Gln	Gla	Lev	Tyr	Cly	Gly
					275					280					285
	Lys	Gin	Gly	Ser	Pro	Рго	årz	Cys	Pro	Leu	Asn	Pro	Gly	Leu	Pro
					290					295					300
35	Pro	Gly	Leu	Leu	Phe	Leu	He	ÅSB	Pro	Lys	Asn	Pro	Thr	Tyr	Gly
				•	305					310					315
	Pro	Asn	He	Cys	Asp	Cly	lsn	Phe	ASP	Thr	Vài	Ala	Hel	Leu.	λrg
40					320					325					330
	Cly	Glu	Het	?he	ASP	Phe	Lys	Lys	Ar:	Trp	Phe	Irp	Ars	Yal	irs
					335					340					345
45	Asn	As'n	Gin	V a I	He t	ASP	Cly	Tyr	Pro	Hei	Pro	He	Gly	Gln	Phe
			:		350					355		•			360
	Trp	Arz	Cly	Leu	Pro	Ala	Ser	He	Àsn	Thr	Ala	lyr	Glu	Arg	Lys
50					206					170					375

	[Sequence Sheet 3]	
•	Sequence No.: 1 (continued)	
5	Asp Gly Lys Phe Val Phe Phe Lys Gly Asp Lys His Trp Val Phi	•
	380 385 390	No.: 1 (continued) 18 Phe Val Phe Phe Lys Gly Asp Lys His Trp Val Phe 380 385 390 18 Ser Leu Glu Pro Gly Tyr Pro Lys His IIe Lys Glu 395 400 405 18 Gly Leu Pro Thr Asp Lys IIe Asp Ala Ala Leu Phe 410 415 420 10 Asn Gly Lys Thr Tyr Phe Phe Arz Gly Asn Lys Tyr 425 430 435 10 Asn Glu Glu Leu Arz Ala Val Asp Ser Glu Tyr Pro 440 445 450 10 Lys Val Trp Glu Gly IIe Pro Glu Ser Pro Arz Gly 455 460 465 11 Gly Ser Asp Glu Val Phe Thr Tyr Phe Tyr Lys Gly 470 475 480 12 Irp Lys Phe Asn Asa Gln Lys Leu Lys Val Glu Pro 485 490 495 18 Pro Asp Glu Gly Thr Glu Glu Glu Thr Glu Val IIe 515 520 525
	Asp Glu Ala Ser Leu Glu Pro Gly Tyr Pro Lys His lie Lys Glo	ı
10	395 400 405	j
	Leu Gly Ars Gly Leu Pro Thr Asp Lys lle Asp Ala Ala Leu Phe	:
	410 415 420)
!5	Trp Net Pro isn Gly Lys Thr Tyr Phe Phe Arz Gly Asn Lys Tyr	
	425 430 435	j
	Tyr Arz Phe Asn Clu Glu Leu Arz Ala Val Asp Ser Glu Tyr Pro	ľ
20	440 445 450)
	Lys Asn lie Lys Val Trp Glu Gly lie Pro Glu Ser Pro Arg Gly	,
	455 460 465	,
?5	Ser Phe Net Gir Ser Asp Giu Val Phe Thr Tyr Phe Tyr Lys Giy	,
	470 475 480)
	Asn Lys Tyr Trp Lys Phe Asn Asa Gin Lys Leu Lys Val Giu Pro)
30	485 490 495	j
	Gly Tyr Pro Lys Ser Ala Leu Ar: Asp. Trp Het Gly Cys Pro Ser	
	500 505 \$10)
35	Gly Gly Arg Pro Asp Clu Gly Thr Glu Glu Glu Thr Glu Yai lle	;
	515 520 525	,
	lle lle Giu Val Asp Ciu Giu Giy Giy Giy Ala Val Ser Ala Aia	l
40	530 535 540)
	Ala Vat Val Leu Pro Val Leu Leu Leu Leu Leu Val Leu Ala Val	
	545 550 555)
45	Gly Leu Ala Yal Phe Phe Phe Arz Arz His Gly Thr Pro Arz Arz	;
	560 565 570)
	Leu Leu Tyr Cys Gin Arg Ser Leu Leu Asp Lys Val	
50 -	575 580	

[Sequence Sheet 4]

Sequence No.: 2

Length of sequence: 3403

Type of sequence: Nucleic acid

Number of chain: Double strand

Topology: Linear

Class of sequence: cDNA to mRNA

Origin:

Species: Human

Tissue: Placenta

	(Sequence She	eet 5]			
5	Sequence No.	2 (continued	i)		
	AGTTCAGTGCCTACC	GAAGACAAAGGCGCC	CCGAGGGAGTGGCGG	TGCGACCCCAGGGCG	60
10	тсссссссссссссс	AGCCACACTGCCCGG	CTGACCCGGTGGTCT	CGGACCATGTCTCCC MetSerPro	120
15	GCCCCAAGACCCTCC AlaProArgProSer	CGTTCTCTCCTCCTC ArgCysLeuLeuLeu	CCCCTGCTCACGCTC ProLeuLeuThrLeu /5	GGCACCGCGCTCGCC GlyThrAlaLeuAla	180
20		CAAAGCAGCAGCTTC GInSerSerSerPhe			240
25		GACCTACGTACCCAC AspLeuArgThrHis 50			300
	GCCATCGCTGCCATG AlalleAlaAlaMet	CAGAAGTTTTACGGC GInLysPheTyrGly 20	TTGCAACTAACAGGC LeuGinVaiThrGly 75	AAAGCTGATGCAGAC LysAlaAspAlaAsp	360
30		AGGCGCCCCCGATGT ArgArgProArgCys		TTTGGGGCTGAGATC PheGlyAlaGlulle	420
35		AGGAAGCGCTACGCC ArgLysArgTyrAla			480
40		CAGAATTACACCCCC GlnAsnTyrThrPro		AlaThrTyrGluAla	540
45		CGCGTGTGGGAGAGT ArgValTrpGluSer			600
-1 0		GAGGGCCATGAGAAG GluGlyHisGluLys 70			660

[Sequence Sheet 6]

Sequence No.: 2 (continued)

	CCCTTCCATGGCGAC	ACGACGCCTTCGAT	GCTGAGGGCGGCTTC	CTGGCCCATGCCTAC	720
	GlyPheHisGlyAsp 185	ArgThrAlaPheAsp /90	GlyGluGlyGlyPhe 195	LeuAlaHisAlaTyr مصد	
0	TTCCCAGGGCCCAAC	ATTGGAGGAÇACACC	CACTTTGACTCTGCC	CAGCCTTGGACTGTC	780
	PheProGlyProAsn ンのS	lleGlyGlyAspThr <i>コル</i>	HisPheAspSerAla ング	GluProTrpThrVal صدد	
_				CACGAGCTGGGCCAT	840
•	ArgAsnGluAspLeu کدد	AsnGlyAsnAsplie のそと	PheLeuValAlaVal	HisGluLeuGlyHis 240	
	GCCCTGGGGCTCGAG	CATTCCAGTGACCCC	TCGCCCATCATGGCA	CCCTTTTACCAGTGG	900
o	AlaLeuGlyLeuGlu	HisSerSerAspPro ユケロ	SerAlalleMetAla	ProPheTyrGInTrp	
	ATGGACACGGAAAAA	TTTGTGCTTCCCCAT	TATGACCCCCGGGGC	ATCCAGCAACTTTAT	960
5	WetAspThrGluLys ککر	PheValLeuProllis ユクロ	TyrAspProArgGly	ileGinGinLeuTyr عد	
9	GGGGGTAAGCAAGGT	TCCCCACCAAGATGC	CCCCTCAACCCAGGA	CTACCTCCCGGCCTT	1020
	GlyGlyLysGlnGly عهد	SerProProArgCys	ProLeuAsnProGly	LeuProProGlyLeu 300	
0				TGTGACGGGAACTTT	1080
	LeuPheLeuIIeAsn 305	ProLysAsnProThr	TyrGlyProAsnile	CysAspG1yAsnPhe シンの	
				CCCTCCTTCTCCCCC	1140
5	AspThrValAlaNet シング	LeuArgGlyGluHet 330	PheaspPheLysLys	ArgTrpPheTrpArg	
				CAGTTCTGGCGGGGC	1200
o	ValArgAsnAsnGln シルケ	ValNetAspGlyTyr 350	ProNetProlleGly	GinPheTrpArgGly	
				TTCGTCTTCTTCAAA	1260
	LeuProAlaSerile 965	AsnThrAlaTyrGlu シクロ	ArgLysAspGlyLys	PheValPhePheLys	٠
5				TACCCCAAGCACATT	1320
	GlyAspLysllisTrp 385	ValPhe∧spGluAla 390	SerLeuGluProGly	TyrProLysHislle	

55

[Sequence Sheet 7]

Sequence No.: 2 (continued)

	AACGAGCTGGGCCGA	GGGCTGCCTACCGAC	AAGATTGATGCTGCT	CTCTTCTGGATGCCC	1380
10		GlyLeuProThrAsp			
		TICTTCCGTGGAAAC PhePheAirgGlyAsn 440		•	1440
15	GCAGTGGATAGCGAG	TACCCCAAGAACATC TyrProLysAsnIle	LysValTrpGluGly	HeProGluSerPro	1500
20		#\$0 GGCAGCGATGAAGTC GlySerAspGluVal			1560
	445	470 CAGAAGCTGAAGGTA	475	480	1620
25	TrpLysPheAsnAsn #85	GInLysLeuLysVal <i>490</i>	GluProGlyTyrPro 495	LysSerAlaLeuArg 500	
		CCATCGGGAGGCCGG ProSerGlyGlyArg			1680
30	GTGATCATCATTGAG	GTGGACGAGGAGGC ValAspGluGluGly			1740
35	525	CTCCTCCTCCTCCTC	क्रद	240	1800
	545	LeuLeuLeuValLeu 550	555	560	
40		AGGCGACTGCTCTAC ArgArgLeuLeuTyr 520			1860
		CCCCCCACTCCTACC		CTGAAGGCCAGTGGC	1920
45	AGCAGGTGGTGGTGG	GTGGGCTGCTCCCAT	CGTCCCGAGCCCCCT	CCCCGCAGCCTCCTT	1980

[Sequence Sheet 8]
Sequence No.: 2 (continued)

10	ссттстстстстссс	стссстссстсстт	CACCCTGACCGCCTC	сстссстсстсссс	2040
	GGCATTGCATCTTCC	CTAGATAGGTCCCCT	GAGGGCTGAGTGGGA	GGGCGGCCCTTTCCA	2100
15	GCCTCTGCCCCTCAG	GGGAACCCTGTAGCT	TTGTGTCTGTCCAGC	CCCATCTGAATGTGT	2160
20	TGGGGGCTCTGCACT	TGAAGGCAGGACCCT	CAGACCTCGCTGGTA	AAGGTCAAATGGGGT	2220
25	CATCTGCTCCTTTTC	CATCCCCTGACATAC	CTTAACCTCTGAACT	CTGACCTCAGGAGGC	2280
	TCTGGGGAACTCCAG	CCCTGAAAGCCCCAG	GTGTACCCAATTGGC	AGCCTCTCACTACTC	2340
30	TTTCTGGCTAAAAGG	AATCTAATCTTGTTG	AGGGTAGAGACCCTG	AGACAGTGTGAGGGG	2400
35	GTGGGGACTGCCAAG	CCACCCTAAGACCTT	GGGAGGAAAACTCAG	AGAGGGTCTTCGTTG	2460
40	CTCAGTCAGTCAAGT	TCCTCGGAGATCTTC	CTCTGCCTCACCTAC	CCCAGGGAACTTCCA	2520
45	AGGAAGGAGCCTGAG	CCACTGGGGACTAAG	TCCCCAGAAGAAACC	CTTGGCAGCCCTGTG	2580
.5	CCTCTCGAATGTTAG	CCTTGGATGGGGCTT	TCACAGTTAGAAGAG	CTGAAACCAGGGGTG	2640

		•	
	[Sequence Sheet 9]		
	Sequence No.: 2 (continued)		
5	CAGCTGTCAGGTAGG GTGGGGCCGGTGGGA GAGGCCCGGGTCAGA	CCCCTGGGGGTGAGC	2700
10	CTTAAGGCCACAGAG AAAGAACCTTGCCCA AACTCAGGCAGCTGG	GGCTGAGGCCCAAAG	2760
15	GCAGAACAGCCAGAG GGGGCAGGAGGGGAC CAAAAAGGAAAATGA	GGACGTGCAGCAGCA	2820
20	TTGGAAGGCTGGGGC CCGGCAGCCAGGTTA AAGCTAACAGGGGGC	CATCAGGGTGGGCTT	2880
	GTGGAGCTCTCAGGA AGGGCCCTGAGGAAG GCACACTTGCTCCTG	ттестссстстсстт	2940
25	GCTGCCCAGGCAGGG TGGAGGGGAAGGGTA GGGCAGCCAGAGAAA	GGAGCAGAGAAGGCA	3000
30	CACAAACGAGGAATG AGGGGCTTCACGAGA GGCCACAGGGCCTGG	CTGGCCACGCTGTCC	3060
35	CGGCCTGCTCACCAT CTCAGTGAGGGACAG GAGCTGGGGCTGCTT	AGGCTGGGTCCACGC	3120
	TTCCCTGGTGCCAGC ACCCCTCAAGCCTGT CTCACCAGTGGCCTG	ссстстссстссссс	3180
40	ACCCAGCCCACCCAT TGAAGTCTCCTTGGG TCCCAAAGGTGGGCA	TGGTACCGGGGACTT	3240
45	GGGAGAGTGAGACCC AGTGGAGGGGAGCGAAG AGGAGAGGGATGTGG		3300
	CTACCGCAAATGGGG TGAACGGTGCTGGCA GTTCGGCTAGATTTC	TETETTETTT	3360

50

55

TITCTTTTCTTIAAT GTATATTTATTAT AATTATTATATAT

[Sequence Sheet 10]

Sequence No.: 3 Length of sequence: 7 Type of sequence: Amino acid Topology: Linear 10 Class of sequence: Peptide Fragment type: Intermediate fragment Sequence 15 Pro Arg Cys Gly Val Pro Asp 5 1 - 20 [Sequence Sheet 11] Sequence No.: 4 25 Length of sequence: 9 Type of sequence: Amino acid 30 Topology: Linear Class of sequence: Peptide Fragment type: Intermediate fragment 35 Sequence Gly Asp Ala His Phe Asp Asp Asp Glu 5 1 40 [Sequence Sheet 12] 45 Sequence No.: 5 Length of sequence: 20 Type of sequence: Nucleic acid 50 Number of chain: Double strand

Topology: Linear

Class of sequence: Other nucleic acid, synthetic DNA

5 Sequence

CC(C/A)(C/A)G(G/A/C)TG(T/C)(C/G)G(G/A/C)(G/A)(A/T)(G/C/T)CC

(T/A) GA

10

20

[Sequence Sheet 13]

Sequence No.: 6

Length of sequence: 25

Type of sequence: Nucleic acid

Number of chain: Double strand

Topology: Linear

25 Class of sequence: Other nucleic acid, synthetic DNA

Sequence

(T/C) TC (G/A) T (G/C) (G/A/C) TC (G/A) TC (G/A) AA (G/A) TG (G/A)

(C/A/T) (G/A) TC (T/C)

35 [Sequence Sheet 14]

Sequence No.: 7

Length of sequence: 27

Type of sequence: Amino acid

Topology: Linear

45 Class of sequence: Peptide

Fragment type: Intermediate fragment

Sequence

50

	Gly	Gly	Gly	Ala	Val	Ser	Ala	Ala	Ala	Val		
·	1				5					10		
5	Val	Leu	Pro	Val	Leu	Leu	Leu	Leu	Leu	Val		
					15					20		
	Leu	Ala	Val	Gly	Leu	Ala	Val	Phe	Phe	Phe		
10					25		•					
. * ,				·								
15	[Seq	uence	Shee	t 15]					٠			
	Sequ	ence	No.:	8								
20	Leng	th of	sequ	ence:	14	,		•				
20	Type of sequence: Amino acid											
•	Торс	logy:	Line	ear			. *					
25	Clas	s of	seque	ence:	Pepti	.de						
	_	ence										
30	Arg	Glu	Val	Pro	Tyr	Ala	Tyr	IIe	Arg	Glu		
•	1				5					10		
	Gly	His	Glu	Lys								
35				•								
	[Sec	quence	e She	et 16	}							
40	Sequ	Sequence No.: 9										
		Length of sequence: 14										
·	Тур	e of	seque	nce:	Amino	acid						
45	=	ology										
	Cla	ss of	sequ	ence:	Pept	ide						
50	Seq	uence										

Asp

Asn

Gly

Asp

Phe

Ala

Val

Met

Leu

1 5 10

Arg Gly Glu Met

[Sequence Sheet 17]
Sequence No.: 10
Length of sequence: 15
Type of sequence: Amino acid
Topology: Linear
Class of sequence: Peptide

Sequence

Gly Trp Met Asp Ala Leu Arg Pro Lys Ser 10 5 1 Gly Gly Pro Ser Cys

15

Claims

10

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- 1. A protein having the amino acid sequence shown in Sequence Sheet sequence number 1.
 - 2. A DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2 which encodes a protein having the amino acid sequence shown in Sequence Sheet sequence number 1.
- 3. A plasmid containing a DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2, and expressing the protein shown in Sequence Sheet sequence number 1.
 - 4. A host cell harbouring a plasmid containing a DNA having the nucleotide sequence shown in Sequence Sheet sequence number 2, and expressing the protein shown in Sequence Sheet sequence number 1.
- Monoclonal antibodies which peculiarly recognize a protein having the amino acid sequence shown in Sequence Sheet sequence number 1.

50 -

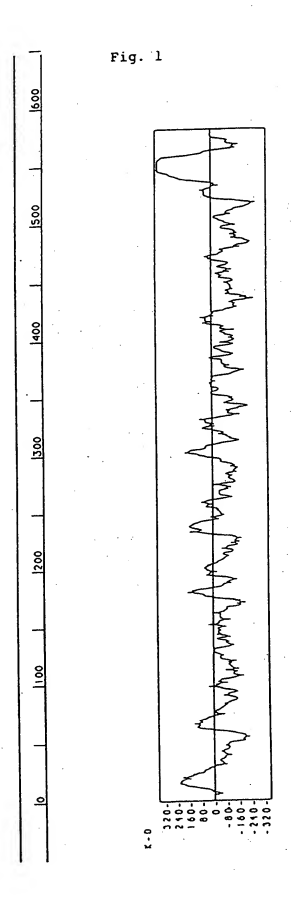


Fig. 2A

MAPAAHLRSA	A AARALLPPML	LLLLQPPPL.	•	· · · · · · LARA	33
MH S	JAGA	LLLLFWGVVS	HSFPATLETQ	EQDVDLVQKY	37
MFSLKTL	PFL	LLLHVÕISKA	FPVSSKEKNT	KTVQD····Ÿ	36
MMHL		VLLCLPVCSA	YPLSGAAKEE	DSNKDLAQQY	37
MKSL		LLLCVAVCSA	YPLDGAARGE	DTSMNLVQKY	37
MS LWQP	LVLÝLLV	LGCCFAAPRQ	RQSTLVLFPG	DLRTNLTDRQ	4 3
•			· · SPIIKFPG	DVAPK·TDKE	19
MR	· ···LTVLCAV	CLLPGSLALP	LPQEAGGMSE	3 MOT	33
MSPAP	RPSR	CLLLPLLTLG	TALASLGSAQ	SSSFS - PEAW	38
Σ				:	50
LP PDVHHL.	·· ···HAERR·G	POPWHAALPS	SP···SPA	TOFAPRPASS	7
L-EKYYNLK	KN DGRQVEKRRN	SGPVVEKLKO	MQEFFGLKVT	GKPDAETLKV	. 8
L-EKFYQLP	PS NOYOSTRKNG	TNVIVEKLKE	MORFFGLNVT	GKPNEETLDM	9 6
L. EKYYNLE	EK DVKQFRRK.D	SNLIVKKIQG	MOKFLGLEVT	GKLDTDTLEV	8 5
L. ENYYDLK	KK DVKQFVRRKD	SGPVVKKIRE	MOKFLGLEVT	GKLDSDTLEV	9.8
LAEEYLYRY	YG YTRVAEMRGE	SKSLGPALLL	LOKOLSLPET	GELDSATLKA	9 6
LAVQYLNTF	F. YGCPKE-SCN	·LFVLKDTLKK	MOKFFGLPQT	GDLDQNTIET	67.
	F. YLYDSETK.N	ANSLEAKLKE	MOKFFGLPIT	GMLNSRVIEI	81
L.QQYGYLP	PP GDLRTHTQRS	PQSLSAAIAA	MQKFYGLQVT	GKADADTMKA	87
L·E.YL.	: : : : : : :	KL	MQKF.GL.VT	GKLDTL	100

MMP - 11	LRPPRCGVPD	PSDGLSARN	ROKRFVLSGG	RWEKTDLTYR	ILRFPWQLVQ	123	
MMP · 1	MKQPRCGVPD	. V AQ - FV L	· · · · TE · GNP	RWEQTHLTYR	IENYTPOLPR	127	
MMP · 8	MKKPRCGVPD	· SGG · FML · ·	TP.GNP	KWERTNLTYR	IRNYTPQLSE	126	
MMP . 10	MRKPRCGVPD	· VGH · FSS · ·	· · · · FP · GMP	KWRKTHLTYR	IVNYTPOLPR	126	
MMP · 3	MRKPRCGVPD	· VGH · FRT · ·	· · · · FP · GIP	KWRKTHLTYR	IVNYTPOLPK	127	
MMP · 9	MRTPRCGVPD	· LGR · FQT · ·	· · · · FE · GDL	KWHHHNITYW	IQNYSEDLPR	134	
MMP · 2 ·	MRKPRCGNPD	· VAN · YNF · ·	· · · · FP · RKP	KWDKNQITYR	IIGYTPDLDP	108	
MMP . 7	MQKPRCGVPD	·VAE·YSL··	FP . NSP	KHTSKVVTYR	IVSYTRDLPH	122	
MT - MMP	MRRPRCGVPD	KFGAEIKANV	RRKRYAIQGL	KWQHNEITFC	IQNYTPKVGE	137	
Consensus	MRKPRCGVPD	. VG . F	F P . G . P	KWT.LTYR	I.NYTPOLP.	Fig.	n: -1
M P · 1 1	EOVROTMAEA	LKVWSDVTPL	TFTEV	· · · · HEGRADI	MIDFARYWDG	2B	20
MMP · 1	ADVDHAIEKA	FOLWSNVTPL	TFTKV	···SEGQADI	MISFVRGDHR	169	
MMP · 8	AEVERAIKDA	FELWSVASPL	IFTRI	···SQGEADI	NIAFYQRDHG	168	
MMP - 10	DAVDSAIEKA	LKVWEEVTPL	TFSRL	···YEGEADI	MISFAVKEHG	168	
MMP · 3	DAVDSAVEKA	LKVWEEVTPL	TFSRL	···YEGEADI	MISFAVREHG	169	
WMP . 9	AVIDDAFARA	FALWSAVTPL	TFTRV	···YSRDADI	VIQFGVAEHG	176	
MMP - 2	ETVDDAFARA	FOVWSDVTPL	RFSRI	···HDGEADI	MINFGRWEHG	150	
MMP - 7	ITVDRLVSKA	LNMWGKEIPL	HFRKV	· · · · VWGTADI	MIGFARGAHG	164	
MT - MMP	YATYEAIRKA	FRVWESATPL	RFREVPYAYI	REGHEKQADI	MIFFAEGFHG	187	
Consensus	VD.AKA	F. VWS. VTPL	TF.RV	···.EG.ADI	MI.FAHG	200	

		_					-	Fig	•	2C			_		١٥	~	~	σ	0
208	-1	210	211	225	199	207	229	250		208	211	210	210	211	275	249	207	229	300
GTD	XX	GTN	GTN	VVVPTRFGNA	OVVRVKYGNA	GIN	D.C.N								DRFGFCPSER	GKYGFCPHEA			
DETWTIGDDQ HERWTNNFT.	EETWTNTSA	DEKWTEDAS.	DEQWT KDTT.	DELWSLGKG.	DELWTLGEG.	DERWTDGSSL	AEPWTVRNE .	DE.WT				•	•	•	CSTTANYDTD	CSTTYNFEKD			
HREGDVHFDY GIGGDAHFDE	GIGGDAHFDA	GLYGDIHFDD	GINGDAHFDD	GIQGDAHFDD	GVGGDSHFDD	GLGGDAHFDE	NIGGDTHFDS	GIGGDAHFD.		•	•	•			TDGRSDGLPW	DTGRSDGFLW			
I LAHAFFPKT NLAHAFQPGP	ILAHAFQPGQ	SLAHAYPPGP	VLAHAYAPGP	LLAHAFPPGP	LLAHAFAPGT	TLAHAFAPGT	FLAHAYFPGP	LAHAF. PGP		•		•	•	•	FEGRSYSACT	FNGKEYNSCT	•		
DDLP FDG PGG DN SP FDG PGG	DNSPFDGPNG	DFYSFDGPGH	DEYPFOGPGN	DGYPFDGKDG	DGYPFDGKDG	DSYPFDGPGN	DRTAFDGEGG	D. YPFDGPGG			•	•		•	DGAACHFPFI	DGEYCKFPFL	•		
MMP - 1 1	MMP · 8	MMP - 10	MMP . 3	WMP · 9	MMP - 2	MMP - 7	MT · MMP	Consensus		MMP - 11	MMP · 1	MMP - 8	MMP - 10	MMP · 3	WMP . 9	MMP · 2	MMP · 7	MM MMP	Consensus

Fig. 2D

MMP - 11			•	•		20.8
MMP - 1		•	•		:	211
MMP · 8			•		•	210
MMP-10						210
MMP - 3	•		•			211
WMP · 9	LYTRDGNADG	KPCQFPFIFQ	GQSYSACTTD	GRSDGYRWCA	TTANYDRDKL	325
MMP . 2	LFTMGGNAEG	QPCKFPFRFQ	GTSYDSCTTE	GRTDGYRWCG	TTEDYDRDKK	299
MMP - 7	•		•	•		207
MT.MMP	•		•			229
Consensus			•			350
MMP - 11		•	•			208
MMP-1		•	:	:	•	211
MMP - 8	•	•	•	•	:	210
MMP-10	•		•	•	:	210
MMP - 3	•		•		:	211
WMP - 9	FGFCPTRADS	TVMGGNSAGE	LCVEPFTFLG	KEYSTCTSEG	RGDGRLHCAT	375
MMP - 2	YGFCPETAMS	TVGG · NSEGA	PCVFPFTFLG	NKYESCTSAG	RSDGKMWCAT	348
MMP - 7		•	•	•	:	207
MT-MMP			•	•		229
Consensus		•		•		400

MMP - 11		J	LQVAA · HEFG	HVLGLQHTTA	AKALMSAFY.	237
MMP - 1	•	· · · · · · · · · · · · · · · · · · ·	HRVAA · HELG	HSLGLSHSTD	IGALMYPSY.	240
MMP · 8			FLVAA · HEFG	HSLGLAHSSD	PGALMYPNY.	239
MMP - 10	•		FLVAA·HELG	HSLGLFHSAN	TEALMYPLYN	240
MMP . 3	•	J	FUVAA·HEIG	HSLGLFHSAN	TEALMYPLYH	241
WMP · 9	TSNFDSDKKW	GFCPDQGYSL	FLVAA - HEFG	HALGLDHSSV	PEALMYPMY.	423
MMP - 2	TANYDDDRKW	GFCPDQGYSL	FLVAA·HEFG	HAMGLEHSQD	PGALMAPIY.	396
MMP - 7			FLYAATHELG	HSLGMGHSSD	PNAVMYPTY.	236
MT-MMP		IOND	FLVAV·HELG	HALGLEHSSD	PSAIMAPFY	261
Consensus			FLVAA·HE.G	HSLGL.HS.D	P. ALMYP.Y.	Fig. O vo
MMP - 11	TE. RYPLSL	SPDDCRGVQH	LYG			2E 8 S C
MMP · 1	TFSGDVQL	AQDDIDGIQA	IYG			261
MMP · 8	AFRETSNYSL	PQDDIDGIQA	IYG			262
MMP - 10	SFTELAQFRL	SQDDVNGIQS	LYG			263
MMP · 3	SLTDLTRFRL	SQDDINGIQS	LYG		•	264
MMP · 9	RF. TEGPPL	HKDDVNGIRH	LYGPRPEPEP	RPPTTTPQP	TAPPTVCPTG	471
MMP · 2	TY TKNFRL	SQDDIKGIQE	LYG	•		417
MMP - 7	GNGDPQNFKL	SQDDIKGIQK	LYGKRSNSRK			267.
MT - MMP	QWMDTEKFVL	PHYDPRGIQQ	LYGGKQGSPP	RCPLNPGLPP	GLLFLINPKN	311
Consensus		squoi.ciq.	LYG		:	200

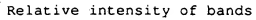
M. O.		M40 · · · · ·	PTVTSRTPAL	GPOAGIDTNE	IAPLEPDAPP	162
MMP · 1	•		RSQNPVQP·I	GPQTP		278
MMP . 8			· · · LSSNP · I	QPTGPST	P KPC	279
MMP - 10	•	O	PPASTEEP.L	VPTKS···VP	S.GSEMPAKC	589
MNIP . 3	•	a. : : :	PPDSPETP.L	VPTEPVP	P. EPGTPANC	290
6. d & &	PPTVHPSERP	TAGPTGPPSA	GPTGPPTA·G	PSTATTV	PLSPVD.DAC	516
MMP . 2		•	ASPDI.D	LGTGPTP	TLGPVTPEIC	440
MMP · 7			•	•		267
MT · MMP	PTYGPNICDG	NFDTVAMLRO	EMFDFKKRWF	HRVRNNQVMD	GYPMPIGQFW	361
Consensus			0.	T.	0	550
		·		•		Fig
MMP - 1.1	DACEASFDAV	STIR-GELFF	FKAGFVWRLR	GGQL · QPGYP	ALASRHWQGL	339
AMP - 1	DS-KLTFDAI	TTIR GEVMF	FKDRFYMR.T	NPFY - PEVEL	NFTSVFWPQL	324 3
. O. X	DP.SLTFDAI	TTLR.GEILF	FKDRYFWR. R	HPQL-QRVEM	NFISLFWPSL	325
MMP · 10	DP - ALSFDAI	STLR-GEYLF		SHWN . PEPEF	HLISAFWPSL	335
MWP - 3	DP - ALSFDAV	STLR.GEILI	FKDRHFWR.K	SLRK·LEPEL	HLISSFWPSL	336
6 · AWW	NV - NI - FDAI		FKDGKYWRFS	EGRGSRPQGP	FLIADKWPAL	563
MMP · 2	KQ.DIVFDGI	AQIR-GEIFF	FKORFIWRTV	TPRO · KPMGP	LLVATFWPEL	487
MMP · 7			•	•		267
MT - MMP	RGLPASINTA	YERKDGKFVF	FKGDKHWVFD	EASLEPGYPK	HIKELGRG · L	410
Consensus	DFDAI	T.R.GEF	FKOR WR		L.S.FWP.L	009

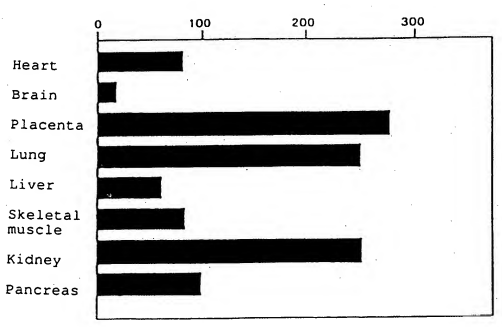
MMP - 11	P.SPVDAAFE	DAQGHIWFF	QG AQYWVYDG	EKPVLG···P	APL-TELGLV	383
MMP · 1	P.NGLEAAYE	FADRDEVRFF	KGNKYWAVQG	QNVLHG · · YP	KDIYSSFGFP	371
MMP · 8	P.TGIQAAYE	DFDROLIFLF	KGNQYWALSG	YDILQGYP	KDI.SNYGFP	371
MMP · 10	P.SYLDAAYE	VNSRDTVFIF	KGNEFWAIRG	NEVQAG - · YP	RGI-HTLGFP	381
MMP · 3	P · SGVDAAYE	VTSKDLVFIF.	KGNQFWAIRG	NEVRAG YP	RGI-HTLGFP	382
W. 9 . 9	P. RKLDSVFE	EPLSKKLFFF	SGRQVWVYTG	ASV-LGP	RRL-DKLGLG	607
MMP · 2	P. EKIDAVYE	APQEEKAVFF	AGNEYWIYSA	STLERG··YP	KPL-TSLGLP	533
MMP · 7		•	•		•	267
MT · MMP	PTDKIDAALF	WMPNGKTYFF	RGNKYYRFNE	ELRAVDSEYP	KNIKVWEGIP	460
Consensus	PDAAYE	(L., (L.,	GN.YWG	GYP	ILG.P	650
						Fi
MMP - 11	R FPVHAAL	VWGPEKNKIY	FFRGRDYWRF	HPSTRRVDSP	VPRRATOWRG	431 ·
MMP · 1	RTVKHIDAAL	S-EENTGKTY	FFVANKYWRY	DEYKRSMDPG	YPKMIAHDFP	4 2 0 0 5 5 6
MMP · 8	SSVQAIDAAV	FYRSKTY	FFVNDQFWRY	DNQRQFMEPG	YPKSISGAFP	4.18
MMP - 10	PTIRKIDAAV	S.DKEKKKŤY	FFAADKYWRF	DENSOSWEOG	FPRLIADDFP	430
MMP · 3	PTVRKIDAAI	S. DKEKNKTY	FFVEDKYWRF	DEKRNSMEPG	FPKQIAEDFP	431
WMP · 9	ADVAQVTGAL	R · SGRGKM · L	LFSGRRLWRF	DVKAQMVDPR	SASEVDRMFP	655
MMP · 2	PDVQRVDAAF	N-WSKNKKTY	IFAGDKFWRY	NEVKKKMDPG	FPKLIADAWN	582
MMP · 7						267
MT · MMP	ESPRGSFM·G	SDEVFTYFYK	GNKYWKFNNO	KLKVEPGYPK	SALRDWMGCP	209
Consensus	. V DAA .	KTY	FF K. WR.	D M. PG	P . I . P	700

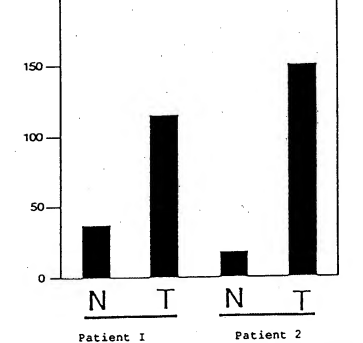
Fig. 2H

796	:		•	C		Consensus
582			DKV·····	RLLYCORSLL	FFFRRHGTPR	MT - MMP
267		•	•		•	MMP · 7
631	•				GSIKSD.WLG	MMP · 2
708		•	•	CPE····DX	···TYD·ILQ	WMP · 9
477				· · · · · · · · · · · · · · · · · · ·	· · · KSNSMLN	MMP · 3
476		•		· · · · · · · · · · · · · · · · · · ·	HTMSNSX···	MMP - 10
468	•			CRY	· · · RGNKWLN	MMP · 8
469	•			CR	KANSWIN	MMP-1
489			NTFLX	CAE	· · · GPD · FFG	MMP - 11
	•		•	•		
750	•	•		٠. >	G D	Consensus
559	LLVLAVGLAV	AAVVLPVLLL	DEEGGGAVSA	EETEVIIIEV	SGGRPDEGTE	MT·MMP
267		•	•	•	•	MMP · 7
621	N-QSLKSVKF	FKGAYYLKLE	·····HS·YF	VV DLQGGG··	AIPDNLDA	MMP · 2
697	QVDQVGYV	WRVSSRSELN	··· CODR · FY	VFQYREKAYF	GVPL··DTHD	WMP · 9
469	AKKVTHTL	GSSQLEFDPN	· · · · · · YF · FT	V. · FEEFGFF	GIDSKIDA	MMP . 3
468	ARMVTHIL··	GSSQFEFDPN	YF.FS	VLQAFGFF	GVEP - · KVDA	MMP - 10
456	AQRVTRVA	GPRYYAFDLI	· · · · · HV · FS	V. · FQQEHFF	GIES··KVDA	MMP . 8
458	TKRILTLQ	GTRQYKFDPK	YF-FH	V··FMKDGFF	GIGH··KVDA	MMP · 1
473	PRLV	PVKVKALEGF	LRGRLYWKFD	FQDADGYAYF	VPSE··IDAA	MMP - 11

Fig. 3







with lung cancer

Relative intensity of Northern blot signals

Patient I

with lung cancer

Fig. 5

MT-MMP cDNA transfection

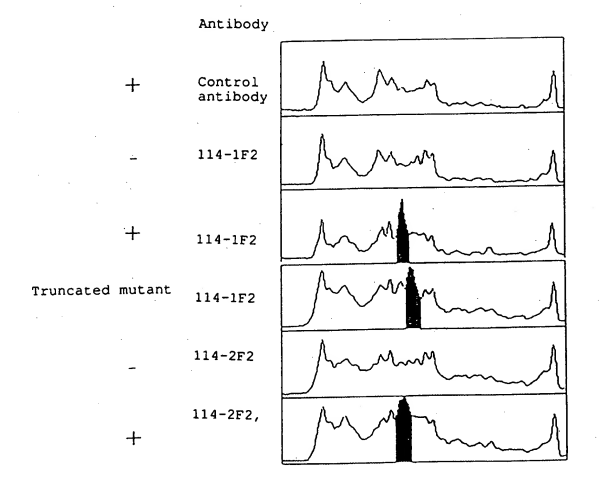


Fig. 6

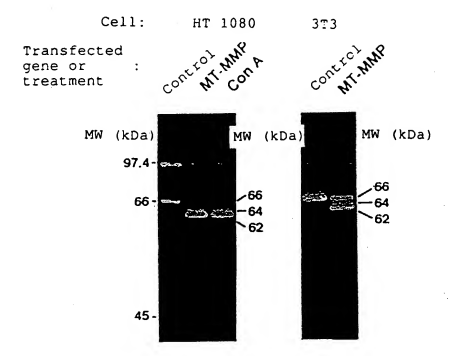


Fig. 7

Cell membrane fraction:

HT 1080 Culture supernatant:

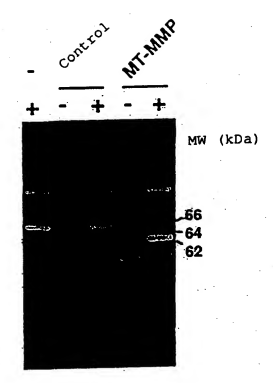
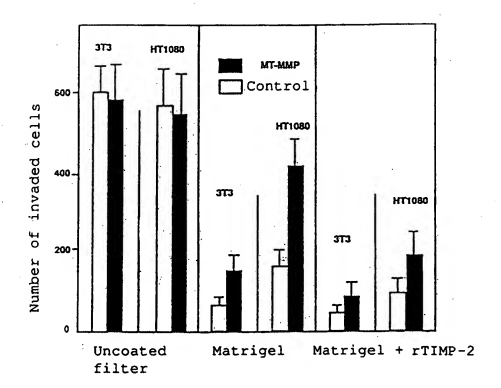


Fig. 8



INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP94/02009

A. CLA	SSIFICATION OF SUBJECT MATTER		
Int.	C16 C12N9/64		
According to	o International Patent Classification (IPC) or to both t	national classification and IPC	
B. FIEL	DS SEARCHED		
	ocumentation searched (classification system followed by	classification symbols)	
	C15 C12N9/50-9/64		
	ion searched other than minimum documentation to the ex		
Electronic da	ta base consulted during the international search (name o	f data base and, where practicable, search to	erms used)
BIOS	IS PREWIEWS, WPI, CAS		
C. DOCU	MENTS CONSIDERED TO BE RELEVANT		,
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
PX	Nature, Vol. 370, No. 6484	·	1-5
	July 7, 1994 (07. 07. 94), Hiroshi Sato et al. "A mat	rix metalloproteinase	
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